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Rigid bronchoscopy in malignant airway fistula using dexmedetomidine

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Introduction

Esophageal and tracheal stenting is an effective palliative treatment of malignant tracheo-esophageal (TEF) and trachea-mediastinal fistula (TMF) to relieve dyspnea and dysphagia (1), and to improve quality of life (2). Rigid bronchoscopy (RB) is commonly used to place tracheal stents to seal the lesion from the airway side (3). However, general anesthesia and jet ventilation required for RB may cause gastric distension, pneumomediastinum or worsening of the fistula (4). Thus, a safe sedation technique enabling RB in spontaneous respiration might be promising.

Dexmedetomidine is an α_2 receptor agonist with sympatholytic and sedative characteristics, which has been shown to enable moderate sedation for several procedures including airway foreign body removal through RB in pediatric patients (5). Apart from one publication (6), to our knowledge there are no reports on tracheal stenting through RB using dexmedetomidine.

Here, we present three cases with successful tracheal stent placement through RB in moderate sedation and spontaneous respiration using dexmedetomidine to seal TEF or TMF. All patients gave the consent to this technique and a general consent to present the data.

Case presentation

Case 1

A 57-year-old male with squamous cell carcinoma (cT4cN2cM0, UICC IIIB) in the posterior mediastinum

presented with an esophageal stent penetrating into the trachea, subsequently causing a 95% stenosis in the lower trachea (*Figure 1A*). Successful placement of a 18-14-14 mm fully covered aerstent[®] Y Carina-Stent (Leufen, Medical GmbH, Bess AG, Berlin, Germany) through a 12mm Storz rigid tracheoscope (Karl Storz SE & Co. KG, Tuttlingen, Germany) was performed to seal the leakage using fluoroscopy (*Figure 1B*). Anesthesia was provided using a continuous infusion pump with remifentanyl (TCI 0.5–1 mcg/mL) and dexmedetomidine. For the latter, a dosage of 4 µg/kg/h over 15 minutes or until Richmond Agitation Sedation Scale (RASS) reached –1 or –2, followed by reduction to 0.7 to 1 µg/kg/h until approximately 10 minutes before end of procedure. In addition, a few propofol boli were added (*Table 1*). The patient tolerated the procedure without negative recalls, was arousable and breathing spontaneously throughout the entire intervention. The postinterventional follow-up was uneventful, and the patient was able to maintain a satisfactory life without dyspnea for 12 months thereafter.

Case 2

A 68-year-old male patient with relapsing small cell lung carcinoma (cT3cN2M0) presented with a 30 mm TMF in the anterior trachea and aspiration pneumonia. TMF was sealed using oxidized regenerated cellulose (Tabotamp[®], Ethicon Inc., Somerville, NJ, USA) and fibrin sealant (Tisseel[®], Baxter AG, Opfikon, Switzerland) (*Figure 2A,B*). Thereafter, an 18-14-14 mm fully covered

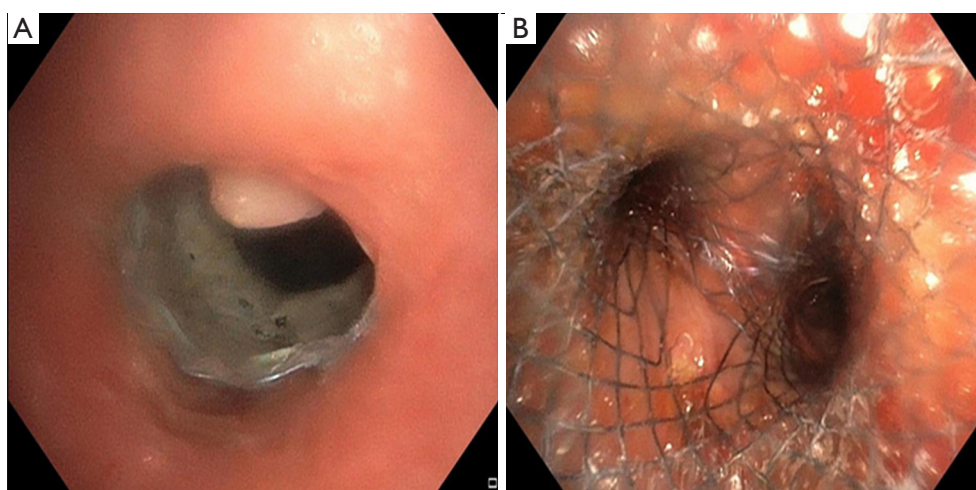


Figure 1 Rigid bronchoscopy of case 1. (A) Upper tracheal view with penetrated esophagus stent causing a 95% stenosis; (B) lower tracheal view after stenting using an aerstent® Y Carina-Stent (Leufen, Medical GmbH, bess AG, Berlin, Germany).

Table 1 Cumulative doses of sedative and analgetic medications for rigid bronchoscopy in spontaneous ventilation

	Case 1	Case 2	Case 3
Propofol (mg)	80	–	120
Remifentanyl (µg)	540	410	490
Dexmedetomidine (µg)	105	150	160

aerstent® Y Carina-Stent was placed through a 12 mm rigid tracheoscope using fluoroscopy (*Figure 2C*). Anesthesia was provided using the identical drug and dose regimen as in case 1 (*Table 1*). The patient remained awake, yet stable and free of dyspnea and pain throughout the whole intervention. The post-interventional follow-up was uneventful. The patient's main symptoms, dyspnea and dysphagia, were relieved. However, the patient died two weeks later of disease progression.

Case 3

A 72-year-old male with relapsing squamous cell carcinoma of the proximal esophagus (cT3cN1M0) presented with an esophageal stent perforating the trachea about 3 cm below the vocal cords and causing 60% tracheal stenosis and complete obstruction of the left main bronchus (*Figure 3A*). Placement of a 16-12-12 mm aerstent® Y Carina Stent with the proximal end right below the cranial fistula orifice (*Figure 3B*) and stent extension

using a partially covered, straight 16×40 mm Ultraflex™ Stent (*Figure 3C*) (BostonScientific, Marlborough, MA, USA) using a 14 mm rigid tracheoscope and fluoroscopy succeeded by administering several propofol boli, remifentanyl and continuous infusion of dexmedetomidine (*Table 1*). A small gap between stent and trachea wall remained at the end (*Figure 3D*), and the esophagus stent was left unchanged. Anesthesia was again provided using a similar drug and dose regimen as in case 1. However, the tapering dose of dexmedetomidine ranged between 1–2.1 µg/kg/h until approximately 10 minutes before end of procedure. The interventions were tolerated without further distress or complications and relieved the patient effectively of his symptoms, most of all dyspnea and dysphagia. Seven days later he died of disease progression.

Discussion

The general condition of patients with TEF/TMF is generally poor due to recurrent aspiration pneumonia and the underlying cancer itself. Additionally, life expectancy precludes major surgery. Endoscopic stenting is the treatment of choice in the majority of cases (1). Eventually immediate closure of the lesion is of highest priority to prevent repeated aspiration. It is recommended to close the esophageal fistula orifice using coated self-expanding metallic stents as initial step (1). For isolated esophageal stenting procedures, analgosedation using combined administration of propofol and opiates seems adequate.

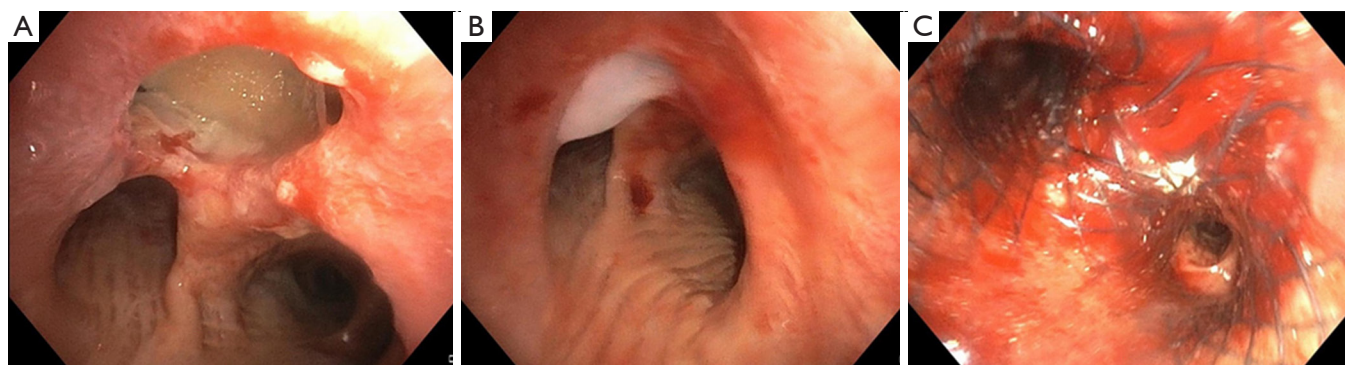


Figure 2 Rigid bronchoscopy of case 2. (A) lower tracheal view revealing a large, supracarinal TMF; (B) lower tracheal view, TMF covered with fibrin sealant; (C) lower tracheal view after stenting using an aerstent® Y Carina-Stent (Leufen, Medical GmbH, bess AG, Berlin, Germany).

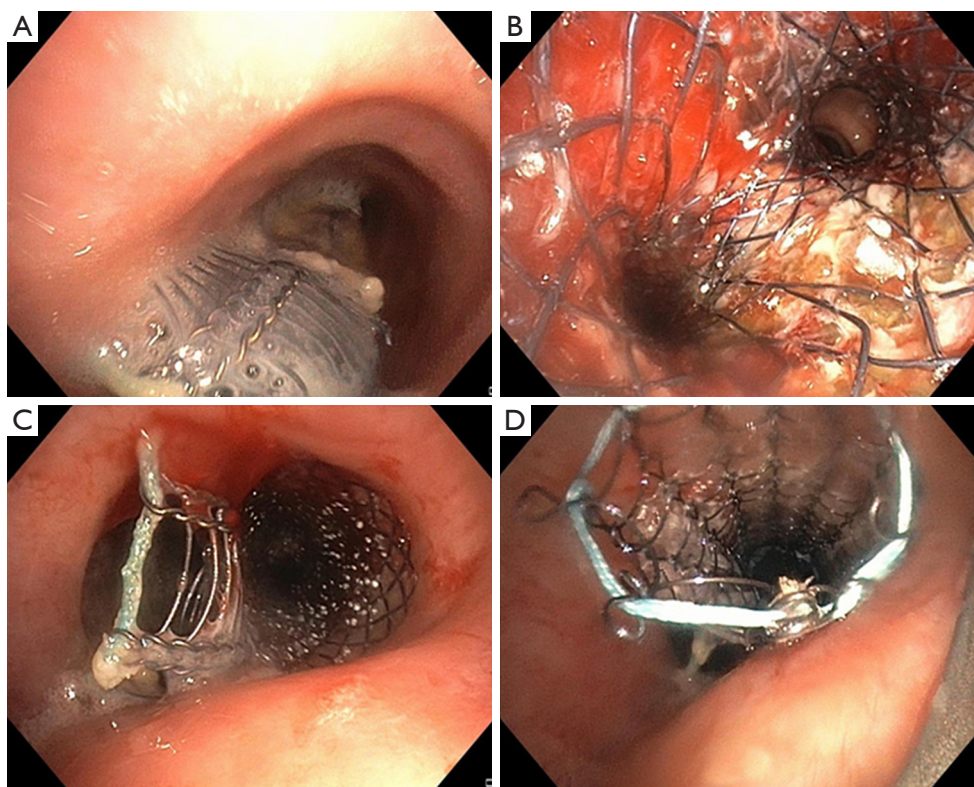


Figure 3 Rigid bronchoscopy of case 3. (A) upper tracheal view with esophageal stent perforating the trachea; (B) lower tracheal view after stenting using an aerstent® Y Carina-Stent (Leufen, Medical GmbH, bess AG, Berlin, Germany); (C) upper tracheal view with persistent TEF above Y stent; (D) showing proximal stent extension using a partially covered, straight 16x40 mm Ultraflex™ Stent (BostonScientific, Marlborough, MA, USA).

However, in case of secondary tracheal compression with symptomatic central airway stenosis or penetration of the esophageal stent into the tracheobronchial system (as shown in case 1 and 3), or in case of TMF (case 2), tracheo-

(bronchial) stenting is the treatment option of choice. In this case, the anesthesiological and airway management becomes more complex. Basically, it is feasible to insert bronchial self-expandable stents using flexible bronchoscopy

in mild sedation only (7). However, TEF/TMF often require insertion of a Y or straight tracheal stent. Rigid bronchoscopy provides ideal prerequisites for safe gas exchange in combination with an adequate working channel for several procedures. To date, tracheal stenting using rigid bronchoscopy unconditionally requires general anesthesia and positive pressure or jet ventilation (1). The latter might need to advance the bronchoscope distally to the fistula, which impedes visualization and intervention. Therefore, the traditional technique requires apnea periods without any ventilation, which further stresses gas exchange and the very limited time to complete stent placement. Eventually, intraoperative fatal outcomes are not uncommon (8). A very recent study by Murgu *et al.* suggests that RB can be safely performed with total intravenous anesthesia and spontaneous ventilation (9). However, in their study dexmedetomidine was only used in two patients without evidence of the procedure details. Our three patients with severe TEF or TMF tolerated RB using dexmedetomidine without pain or stress after careful local anesthesia using lidocaine 10% spray. During potentially painful or stressful periods, its effect was supplemented with remifentanyl and propofol. In addition, dexmedetomidine has been used in combination with small bolus' of propofol (10–20 mg) to suppress coughing and for a safe insertion of the rigid bronchoscope according to Kim *et al.* (10). The sedation has been estimated by clinical assessment and the RASS (Richmond Agitation and Sedation Scale) with the aim of –1 to –2. Spontaneous breathing with sufficient (pre-) oxygenation was maintained at all time in each patient using a face mask and through the bronchoscope, respectively. Meanwhile, the patients were monitored with non-invasive oxygen saturation, ECG, and arterial line to measure blood pressure and arterial O₂ and CO₂.

Eventually, a consented back-up plan in case of under- or oversedation must be considered before starting the procedure (i.e., return to traditional technique with general anesthesia and jet ventilation). Relevant prerequisites for this technique is ensuring that anatomy allows for intubation with a rigid bronchoscopy (oral opening, reclinability) and a trustful patient-doctor relationship. Furthermore, the patient must be informed that he will not be able to speak during the procedure. Thus, a clear hand signaling must be arranged.

In conclusion, the use of dexmedetomidine was of great benefit for the patients and confirmed the results of previous studies regarding spontaneous respiration and moderate sedation. Based on our experience, using dexmedetomidine

spontaneous respiration can be safely maintained even during RB for interventional repair of TEF/TMF.

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Footnote

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Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at <http://dx.doi.org/10.21037/jtd-20-1780>). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. All participants gave informed consent before taking part of this case series.

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